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                 MEDLINE and LMEDLINE reloaded
        MAR 03
NEWS
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        MAR 03
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                 FRANCEPAT now available on STN
        MAR 03
NEWS
      8
                 Pharmaceutical Substances (PS) now available on STN
NEWS
     9
        MAR 29
                 WPIFV now available on STN
        MAR 29
NEWS 10
                 New monthly current-awareness alert (SDI) frequency in RAPRA
         MAR 29
NEWS 11
                 PROMT: New display field available
NEWS 12
         APR 26
                 IFIPAT/IFIUDB/IFICDB: New super search and display field
NEWS 13
        APR 26
                 available
                 LITALERT now available on STN
NEWS 14
         APR 26
                 NLDB: New search and display fields available
NEWS 15
         APR 27
                 PROUSDDR now available on STN
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                 EXTEND option available in structure searching
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                 Polymer links for the POLYLINK command completed in REGISTRY
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         May 12
NEWS 20
        May 17
                 FRFULL now available on STN
                 STN User Update to be held June 7 and June 8 at the SLA 2004
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                 New UPM (Update Code Maximum) field for more efficient patent
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         May 27
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                 CAplus super roles and document types searchable in REGISTRY
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         May 27
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        May 27
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NEWS EXPRESS
              MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 26 APRIL 2004
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NEWS WWW
              CAS World Wide Web Site (general information)
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19 ANSWERS

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L3 15 L2

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L3 ANSWER 1 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 2003:589504 CAPLUS
DOCUMENT NUMBER: 139:13336f
INVENTOR(S): Bertrand, Guyr komanenko, Vadim D.; Raynier, Bernard;
Derrieu, Guy
Vithac S.A., Fr.
SOURCE: EU.; Pat. Appl., 18 pp.
CODEN: EEXENDW
DOCUMENT TYPE: COEN: EXENDW
LARGUAGE: Endish

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: English

				KI	ND					PPLI				DATE			
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L.F	1331	DT.	BE.	CH.	DE.	DK.	ES.	FR.	GB.	GR.	IT.	LI.	LU.	NL,	SE,	MC,	PT,
	г.	TE.	ST.	LT.	LV.	FI,	RO.	MK.	CY.	AL.	TR						
wo	2003	0643	84	Ā	2 . ,	2003	0807	,	W	20	03-E	P151	5	2003	0128		
	2003																
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	ΒY,	BZ,	CA,	CH,	CN,
		co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	ΜX,	ΜZ,	NO,	ΝZ,	OM,	PH,
														TN,			
		UA,	UG,	US,	UΖ,	VC,	٧N,	YU,	ZA,	ZM,	zw,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,
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	RW:	GH,	GM,	KE,	LS,	MW,	ΜZ,	SD,	SL,	SZ,	ΤZ,	UG,	zm,	ZW,	ΑT,	BE,	ВG,
														ΙE,			
		NL,	PT,	SE,	SI,	SK,	TR,	BF,	ΒJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GΩ,	GW,
		ML,	MR,	NE,	sn,	TD,	TG										
														2002			

NIL, PT, SE, St, SN, TW, BE, BD, CF, GG, GF, GN, GN, GW, GW, MD, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO.:

OTHER SOURCE(S):

MARPAT 139:133561

AB RICO(RZCO)NS(O)R [RIR2 = optionally substituted or annelated C1-C20, linear, branched or cyclic alkanediyl, alkenediyl, alkynediyl; R = (un)substituted alkyl) were prepd. for use as haloalkylsulfinylating agents. Thus, lithiosuccinimide was treated with F3CS(O)Cl to give N-trifluoromethylsulfinylsuccinimide which was treated with 1-phenyl-3-methyl-5-aminopyrazole too give the 4-trifluoromethylsulfinylderiv. in 821 yield.

IT 569337-28-6F, 1-(2,4,6-Trichlorophenyl)-3-cyano-4-trifluoromethylsulfinyl-5-aminopyrazole

RL: SPN (Synthetic preparation): PREP (Preparation) (prepn. of trifluoromethylsulfinylsuccinimide as trifluoromethylsulfinylsuccinimide as trifluoromethylsulfinylating agent)

RN 569337-28-6 CAPPUS

CN 1H-Pyrazole-3-carbonitrile, 5-amino-1-(2,4,6-trichlorophenyl)-4-[(trifluoromethylsulfinyl]- (9CI) (CA INDEX NAME)

ANSWER 2 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN

CESSION NUMBER:

ACCESSION NUMBER:

Method for preparation of new N-substituted derivatives of 5-amino-1-phenylpyrazole, the derivatives, and their use as paramiticidal and/or insecticidal agents
Bertrand, Guy; Romanenko, Vadim D.; Raynier, Bernard;
Derrieu, Guy
Virbac SA, Fr.
Fr. Demande, 87 pp.
CODEN: FRXXBL
Patent
French
1

INVENTOR (5):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

FR 2834288 Al 20030704
PRIORITY APPIN. INFO::
CASRFBOW. APPLICATION NO. DATE Al 20030704 FR 2001-17018 20011228 FR 2001-17018 20011228 CASREACT 139:85341; MARPAT 139:85341

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The invention provides new derivs. of 5-amino-1-phenylpyrazoles,
specifically I [wherein: A, B = H, (a)cyclic alk(en/yn)yl (optionally
substituted by one or more halo, alkoxy, alkylthio, or alkoxycathonyl),
halo, cyano, thiocyano, nitro, sulfamido, (di)alkylamino, aminocarbonyl,
aminothiocarbonyl, (di)alkylaminocarbonyl, (di)alkylaminothiocarbonyl,
alkylcarbonylamino, alkylthiocarbonylamino, S(0)nR [n = 0, 1, or 2, and R
= (a)cyclic, (un)satd. (halo)alkyl), Ph, phenylalkyl, or 4- to 7-membered
heterocycly with 1-3 N/o/S/Si atom(m); R1, R2, R3, R4, and R5 = H, halo,
(a)cyclic (un)satd. C1-6 (halo)alkyl, (halo)alkoxy, or (halo)alkylthio; Z
= -N:C:0, -N:C:5, -N:S:0, -NHC(:X)R6, -NHC(:X)XR6, -NHC(:S)XR6,
-NHC(:X)RXR8; X = 0 or S; R6 = (un)aubstituted (a)cyclic alk(en/yn)yl,
Ph, phenylalkyl, or heterocyclyl; R7, R8 = H, groups given for R6,
ric

Ph. phenylaikyl, or heterocyclyl; R7, R8 = H, groups given for R6, dimeric unit of I; also Z = -N:C:N- forming a dimer of I; or Z = (un|substituted 1,2-th|azin-2-y| 1-oxide motif). The invention also comprises processes for prepn. of I from corresponding amines I [Z = NH2], typically via reaction of the amines with phosgene, thiophosgene, or thionyl chloride, and optionally reaction of the resultant I [Z = isocyanato, isothlocyanato, or N-sulfinylamino (i.e., -N:S:O]]. Compds. I can be administered to vertebrates, particularly domesticated animals, either orally, topically, or parenterally. In general, I can be used to control both arthropods and nematodes which are parasites of both animals and plants, by application to either the hosts or their environments. Over

specific compds, were claimed per se. Examples (23) include synthesis, and both agrochem, and pharmaceutical formulations. For instance, the amine precursor II [z=Nk2] reacted with phospens in anhyd. PhNe in the presence of 2 equiv pyridine to give II [z=isocyanato] in 95% yield. Reaction of this isocyanate with 3,5-bis(trifluoromethyl)aniline gave title compd. III. Compds. I were against the stablefly Stomoxys calcitrans in a Petri dish expt., at dosages of 0.1 to 30 .mu.g/fly. An

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ANSWER 1 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L3 ANSWER 2 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued) exemplary injectable contained 1% I, 30% Et oleate, and sesame oil qsp 100%, and was sterilized by membrane filtration.

IT \$54415-74-67, 1-(2,4,6-Trichlorophenyi)-3-cyano-4 (trifluoromethyi)thioj-5-(N-sulfinylaminojpyrazole RL: AGR (Agricultural use); IMF (Industrial manufacture); PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOI (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (drug candidate; prepn. of aminophenylpyrazole derivs. as parasiticides

parasiticides
and insecticides)
RN 554415-74-6 CAPLUS
CN 1K-Pyrazole-3-carbonitrile,
5-(sulfinylamino)-1-(2,4,6-trichlorophenyl)-4[(trifluoromethyl)thio]- (9CI) (CA INDEX NAME)

IT 554415-70-2P, 1-(2,4,6-Trichlorophenyl)-3-cyano-4[(trifluoromethyl)sulfinyl)-5-isocyanatopyrazole 554415-71-3P
RL: AGR (Agricultural use): IMF (Industrial manufacture): PAC
(Pharmacological activity): SPN (Synthetic preparation): TRU (Therapeutic
use): BIOL (Biological study): PREP (Preparation): USES (Uses)
(drug candidate): prepn. of aminophenylpyrazole derivs. as
parasiticides
and insecticides!

siticides
and insecticides)
554415-70-2 CAPLUS
1H-Pyrazole-3-carbonitrile, 5-isocyanato-1-(2,4,6-trichlorophenyl)-4[(trifluoromethyl)sulfinyl)- (9CI) (CA INDEX NAME)

NCO

554415-71-3 CAPLUS 1H-Pyrazole-3-carbonitrile, 5-isocyanato-1-(2,4,6-trichlorophenyl)-4-(trifluoromethyl)thio]- (9CI) (CA INDEX NAME)

L3 ANSWER 2 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE

L3 ANSWER 3 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 2002:107926 CAPLUS
DOCUMENT NUMBER: 136:146526
TITLE: Preparation of 1-phenylpyrazole derivatives as insecticides
INVENTOR(S): Manning, David Treadway; Pilato, Michael; Wu, INVENTOR(S): Tai-Teh; Hawkins, David William Rhone-Poulenc Agrochimie, Fr. U.S. Pat. Appl. Publ., 17 pp. CODEN: USXXCO Patent English PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. PATENT NO. KIND DATE US 2002016468
PRIORITY APPLN. INFO.:
OTHER SOURCE(S): A1 20020207

The 1-arylpyrazole-3-thiocarboxamide derivs. I (R1 = H or halo; R2, R4 = R1 or alkyl; R3 = halo, haloalkyl, haloalkoxy or R650m; R5 = alkyl, haloalkyl, alkenyl, alkynyl or cycloalkyl; R6 = alkyl or haloalkyl; Z =

halo, alkyl, formyl, etc.; M = N, C-halo, C-Me, etc.; m, n = 0, 1 or 2) are prepd. as insecticides, esp. active against sucking insects. 146628-02-69 IT

146628-02-69
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (intermediate in prepn. of pyrazolethiocarboxamide insecticide) 146628-02-6 CAPLUS
H-Pyrazole-3-carbonitrile, 5-amino-4-[(dichlorofluoromethyl)thio]-1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

L3 ANSWER 3 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

L3 ANSWER 4 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 1998:479512 CAPLUS
129:95489
TITLE: Preparation of pyridylpyrazole derivatives as pesticides
INVENTOR(S): Manning, David Treadway; Pilato, Michael; Wu, INVENTOR (S): Tai-teh; PATENT ASSIGNEE(S):

Hawkins, David William Rhone-Poulenc Agrochimie, Fr.; Wu, Tai-Teh; Hawkins, David William PCT Int. Appl., 52 pp. COODN: PIXXD2 Patent English DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

20000131 MX 1999-5963 19990623 20001130 BG 1999-103592 19990719 WS 1996-33885P P 19961224 WO 1997-EP7116 W 19971218 CASREACT 129:95489; MARPAT 129:95489 OTHER SOURCE(S):

CSNH2

L3 ANSWER 4 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN

AB The title compds. [I; W = C(:S)NH2; Rl = H, halo; R2, R4 = H, halo, alkyl;

R3 = halo, haloalkyl, etc.; R5 = lower alkyl, haloalkyl, alkenyl, etc.; Z
= H, halo, alkyl, etc.; R = 0-2; M = C-halo, C-Me, N, etc.] are prepd. I
are useful as pesticides. Thus, I (W = CN, R1 = Cl, R2 = R4 = H, R3 = CF1, R5 = Me, Z = H, N = 0, M = C-Cl) was treated with 15-crown-5 in DMF and aq. sodium hydrosulfide to give the title compd. I (W = C(:S)NH2, R1-R5, Z, N, M = same as above), which showed activity against Aphis gossypii and Schizaphis graminum at 10 ppm.

IT 146528-02-6P
RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOI (Biological study); PREP (Preparation); USES (Uses) (prepn. of pyridylpyrazole derivs. as pesticides)

RN 146628-02-6 CAPLUS
CN 1H-Pyrazole-3-carbonitrile, 5-amino-4-[(dichlorofluoromethyl)thio]-1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME) The title compds. [I; W = C(:S)NH2; R1 = H, halo; R2, R4 = H, halo,

REFERENCE COUNT:

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 5 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued) trifluoromethylthiopyrazole with 4-trifluoromethylphenylboronic acid in the presence of PdZ(dba)3, RZCO3 in diglyme afforded I (X = N; Y = C(CN); W = C(SCF3); R5 = NH2; Z = C(C1); R12 = C1; R13, R15, R22, R23, R25, R26

H; R24 = CF3]. The prepd. compds. I showed rather good activity on C. elegans.
207136-58-1
RE: RCT (Reactant); RACT (Reactant or reagent)
(prepn. of pesticidal 1-polyarylpyrazoles)
207136-58-1 CAPINS
HP-Pyrazole-3-carbonitrile, 5-amino-1-(4-bromo-2,6-dichlorophenyl)-4[(trifluoromethyl)thio]- (9CI) (CA INDEX NAME)

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

L3 ANSWER 5 OF 15
ACCESSION NUMBER:
DOCUMENT NUMBER:
1998/304135 CAPLUS
128:321643
Freparation of pesticidal 1-polyarylpyrazoles
Freparation of pesticidal 1-polyarylpyrazoles DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: US 2002002195 US 6433002 20020813 20021210 US 37936 US 2002193411 US 6608093 E Al B2 US 2001-903990 US 2002-152806 20010713 20020523 20021219 20030819 US 1996-30128P US 1997-963631 US 1998-216878 US 2000-606185 PRIORITY APPLN. INFO.: A3 19971104 A3 19981221 A3 20000629 A3 20010412 US 2000-606185 US 2001-832861 MARPAT 128:321643 OTHER SOURCE(S):

The title compds. [I; X = N, CR2; Y = N, CR3; W = N, CR4; R2, R3 = H, halo, OH, etc.; R4 = H, halo, alkyl, etc.; R5 = H, halo, CHO, etc.; Z = H

CR16; R12, R13, R15, R16 = H, halo, alkyl, etc.; R22-R26 = halo, alkyl, haloalkyl, etc.], useful to control pests, were prepd. Thus, reaction of 5-amino-3-cyano-1-(2,6-dichloro-4-bromophenyl)-4-

L3 ANSWER 6 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1996:590912 CAPLUS
125:240887
Pesticidal 1-aryl-5-(substituted alkyl (thio) amidol pyrazoles
Huang, Jamín; Phillips, Jennifer L.
Rhone-Poulenc Inc., USA
U.S., 24 pp., Cont.-in-part of U.S. Ser. No. 21, 717, abandoned.

CODEN: USXXAM Patent

DOCUMENT TYPE: English

LANGUAGE: E.
FAMILY ACC, NUM. COUNT: 1
PATENT INFORMATION:

US 5556873 A 19960917 US 1993-169944 19931220 EP 811615 A1 19971210 EP 1996-108831 199600603 R: AT, BE, CH, DE, DK, ES, FR, GR, GR, IT, LI, LU, NL, NL, SE, PT, IE PRIORITY APPLN. INFO::

US 1993-21717 19930224 US 1993-169944 19931220 OTHER SOURCE(S): MARPAT 125:240887

Novel 1-aryl-5-(substituted alkyl (thio)amido)pyrazoles wherein preferred compds. are of the formula (I) wherein: R is R6S(O)n in which n is 0, 1

2 and R6 is alkyl, preferably methyl; or haloalkyl, preferably trihalomethyl or dihalomethyl; and in which halo is F, Cl or Br or combinations thereof and most preferably CF3, CCl3, CF2Cl, CFCl2, CF2Br, CHF2, CHClF or CHCl2; R1 is H or alkyl; R2 is H or alkyl; R1 and R2

be together to form a 3-7 membered cyclic ring system: R3 is alkoxy, alkoxy(alkoxy)b [b=1-2], alkoxy(alkoxy)b alkyl [b=0-2], alkyls(0)c (c=0 1, 2], alkyls(0)c (c=0)r, phenylalkoxy, pyridylcoxy, pyridyl-5(0)c, optionally substituted with alkyl, halogen, alkoxy, haloalkyl, haloalkoxy, nitro, cyano, alkylthic. R2 and R3 could be together to form a 4-7 membered cyclic ring with 1-2 heteroatoms (e.g. O, S, S(0), S(0)2, NH, N-alkyl); R4 is: hydrogen;

l, preferably methyl; or halogen, preferably F, Cl or Br; R5 is: halogen, preferably F, Cl or Br; alkyl. preferably methyl; haloalkyl, preferably trihalomethyl and more preferably trifluoromethyl; or haloalkoxy, preferably trifluoromethyl; or haloalkoxy, and in

ANSWER 6 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued) which halo is F, Cl or Br or combinations thereof; and X is a nitrogen atom or C-R7 in which R7 is: hydrogen; halogen, preferably F, Cl or Br; cyano; alkyl, preferably Me or ethyl; alkylthio, preferably methylthio or ethylthio; or alkoxy, preferably methylthio or ethylthic or alkoxy, preferably methylthio or ethylthic or insecticides.

IT 181814-41-5P
RL: BaC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation): RIOL (Biological study, unclassified); SPN (Synthetic preparation): RIOL (Biological study)

logical study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (preph. and pesticidal activity of 1-aryl-5-(substituted alkyl (thio) amido)pyrazoles) 181814-41-5 CAPLUS Acctamide, N-[3-cyano-1-(3,5-dichloro-2-pyridinyl)-4-[(trifluoromethyl)thio]-1H-pyrazol-5-yl]-2-methoxy- (9CI) (CA INDEX

ANSWER 7 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

The invention describes novel 1-aryl-5-(substituted alkylideneimino)pyrazole of formula (I) wherein typically preferred substituents are: R1 is cyano, nitro, or halogen; R2 is R9S(O)n in which

n

is 0, 1 or 2 and R9 is alkyl, preferably Me which is substituted by halogen atoms which are the same or different up to full substitution of the alkyl moiety; R3 is hydrogen or alkyl; R4 is Ph or heteroaryl, optionally substituted by one or more hydroxy, halogen, alkoxy, alkylthio,

cyano or alkyl or combinations thereof; preferably R4 is Ph, which is at least substituted by 3-hydroxy or 4-hydroxy; R5 is hydrogen, alkyl or halogen; R6 and R8 are hydrogen; R7 is halogen, alkyl, haloalkyl or haloalkoxy; and X is a nitrogen atom or CR14 in which R14 is hydrogen, halogen, cyano, alkyl, alkylthio or alkoxy. The invention further describes processes to make the compds., compns. of the compds., and methods of use of the compds. for the control of arthropods (mites, aphids

aphids
or insects), nematodes, helminths, or protozoa. Pesticidal activity of I compds. providing 70-100% pest mortality was evaluated against buckthorn aphid, cotton aphid, southern armyworm, Mexican bean beetle, housefly, tobacco budworm, southern corn rootworm, western corn rootworm.

IT 145768-03-2P
RL: BRC (Biological activity or effector, except adverse); BSU
(Biological style unpolassified): SDN (Suppletio preparation): ETC (Piological activity or effector)

logical study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PRDP (Preparation) (pesticidal 1-aryl-5-(substituted alkylideneimino)pyrazoles) 145768-03-2 CAPLUS 1.45768-03-2 CAPLUS 1.4-Pyrazole-3-carbonitrile, -bromo-2,6-dichlorophenyl)-5-[((4-hydroxy-3-methoxynphenyl)methylene)amino]-4-[(trifluoromethyl)thio]- (9CI) (CA INDEX NAME)

L3 ANSWER 7 OF 15
ACCESSION NUMBER:
DOCUMENT NUMBER:
1295:229475 CAPLUS
122:239694
Pesticidal 1-aryl-5-(substituted alkylideneimnio) pyrazoles
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:
U.S., 24 pp. Cont.-in-part of U.S. Ser. No. 790,449, abandoned. abandoned. CODEN: USXXAM DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. DATE KIND DATE PATENT NO. 19941101 19930817 19921031 19921105 19941201 19960331 19921102 US 1992-842431 US 1991-693580 CA 1992-2067282 AU 1992-15192 19920304 19910430 19920427 19920427 US 5360910 US 5236938 US 5236938 CA 2067282 AU 9215192 AU 655014 IL 101702 NO 9201639 EP 511845 EP 511845 IL 1992-101702 19920427 A Al Bl NO 1992-1639 19920428 EP 1992-303857 19920429 19921104 20011031 B1 20011031
CH, DEP, DK, ES, FR, GB, GR, IT, LI, LU, NL, PT, SE
A2 1930128
B1 19970828
B1 19970829
B1 19970827
CI 19970827
RU 1992-294383 19920429
CI 19970827
RU 1992-5011630 19920429
T3 20020316
ES 1992-303857 19920429
T 20020429
PT 1992-303857 19920429
T 20020429
PT 1992-303857 19920429
A 19921124
CN 1992-303857 19920430
B 20000621
A 19921124
BR 1992-1735 19920430
A 19930127
A 19930127
BR 1992-3175 19920430
A 19930127
D 1992-3175 19920430
A 19930615
D 19920430 EP 511845
R: AT, BE,
HU 61529
HV 213630
HV 213630
FV 169737
RU 2088576
AT 207904
ES 2165353
CN 1066265
CN 1063659
BR 9201735
ZA 9203175
JP 05148240
JP 3248943
RO 107407
KC 279252 E T3 T A B A A A B2 B1 BR 1992-1735 2A 1992-3175 JP 1992-111958 19930615 19920430 20020121 19931130 RO 1992-598 19920430 19920430 19920430 19920430 A2 19910430 B2 19911112 A 19920304 A 19920430 SK 1992-1337 CZ 1992-1337 US 1991-693580 US 1991-790449 US 1992-842431 SK 279252 В6 19980805 CZ 286232 В6 20000216 PRIORITY APPLN. INFO .: cs 1992-1337

OTHER SOURCE(S): MARPAT 122:239694

ANSWER 7 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

L3 ANSWER 8 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 1994:527876 CAPLUS
DOCUMENT NUMBER: 121:127876
TITLE: Pesticidal 1-aryl-5-(substituted

n-cinnamylideneimino)

INVENTOR(S): PATENT ASSIGNEE(S):

SOURCE: abandoned.

pyrazoles Huang, Jamin; Manning, David T. Rhone-Poulenc Inc., USA U.S., 21 pp. Cont. of U.S. Ser. No. 71,163,

CODEN: USXXAM COPEN: L
Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

US 5321040 A 19940614
PRIORITY APPLIN. INFO:
OTHER SOURCE(S): APPLICATION NO. DATE A 19940614 US 1993-144262 19931028 US 1993-71163 19930602 CASREACT 121:127876; MARPAT 121:127876

$$R^{1}$$
 N
 $N = CR^{3}CR^{9} = CR^{4}R^{10}$
 R^{5}
 R^{6}
 R^{8}

Novel l-aryl-5-(substituted alkylideneimino)pyrazoles (I; Rl = cyano, nitro or halogen; R2 = unsubstituted or substituted RllS(0)n, in which n AB

0, 1 or 2 and R11 = alkyl or haloalkyl; R3 = R9 = R10 = H; R4 = unsubstituted or substituted Ph or pyridyl; R5 = H, halogen or alkyl; R6

R8 = H or F; R7 = halogen, alkyl, haloalkyl or haloalkoxy; X = N or C-R16 in which R16 = H, halogen, CN, alkyl, alkylthio or alkoxy) were prepd.

used for the control of arthropods (mites, aphids or insects), nematodes, helminths, or protozoa.
157043-81-79

187043-81-78
RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of, as pesticide)
187043-81-7 CAPLUS
HI-Pyrazole-3-carbonitrile, 1-(4-bromo-2,6-dichlorophenyl)-5-[[3-(3-hydroxy-4-methoxyphenyl)-2-propenylidene]amino]-4-

L3 ANSWER 9 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
120:298625 Preparation of phenylpyrazoles as arthropodicides,
nematocides, protozoacides, and anthelmintics
Hatton, Leslie R.; Buntain, Ian G.; Hawkins, David INVENTOR(S):

DOCUMENT TYPE:

Parnell, Edgar W.; Pearson, Christopher J.

PATENT ASSIGNEE(S): SOURCE:

UK.U.S., 76 pp. Cont.-in-part of U.S. Ser. No. 445,153, abandoned.
CODEN: USXXXMP
Patent
English
4

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

							DATE
PATENT NO.	KIND	DATE	:		APPLICATION N	٠.	DATE
					us 1990-52029		19900507
US 5232940	A		0803		TL 1988-86493	,	19880525
IL 86493	A1		1115				
IL 105138	A1		0826		IL 1988-10513	3	19880525
HU 210668	В		0628		HU 1991-1577		19880610
US 5547974	A		0820		US 1993-57669		19930505
FI 9501839	A		0418		FI 1995-1839		19950418
US 5608077	A		70304		US 1995-45441		19950530
US 5714191	A		10203		US 1995-45308		19950530
US 5916618	A	1999	0629		US 1997-94705		19971007
US 6372774	B1		20416		US 1999-35490	3	19990716
DK 200201527	A5	2002	21010		DK 2002-1527		20021010
PRIORITY APPLN. INFO.	:			GB	1985-31485	А	19851220
					1986-943132	B1	
					1987-13768	А	19870612
					1987-13769	Α	19870612
				US	1988-205238		19880610
				บร	1988-205299	В1	19880610
				US	1989-380333	В1	19890717
				US	1989-413134	В1	19890927
				US	1989-445153	B2	
				IL	1986-81025	А	19861218
				IL	1988-86492	Α	19880525
				DK	1988-3140	L	19880609
				FI	1988-2735	А	
				НU	1988-3009	A	19880610
				US	1990-520290	A3	19900507
				ŲS	1993-57669	A3	19930505
				US	1995-453087	A1	19950530
				US	1996-652921	В1	19960524
				US	1997-855876	В3	19970512
				US	1998-137313	В3	19980821
OTHER SOURCE(S):	MA	RPAT	120:298	625			

ANSWER 8 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN [(trifluoromethyl)sulfinyl]- (9CI) (CA INDEX NAME)

(Continued)

L3 ANSWER 9 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN

Title compds. [I; R1 = cyano, nitro, halo, acetyl, formyl, (halo)alkyl, etc.; R2 = R'502, R'50, R'S, halo, cyano, nitro, cycloalkyl, alkenyl, thiocyanato, sulfamoyl, carbamoyl, alkoxycarbonyl, alkanoyl, (halo)alkyl; R' = (substituted) alkyl, alkenyl, alkynyl; R3 = H, (substituted) amino, alkoxycarbonyl, alkoxymethyleneamino, halo, cycloalkyl, cycloalkylcarbonyl, alkylsulfenylamino, trialkylsilylmethyl, etc.; R4-R8

H, halo, nitro, cyano, (halo-substituted) alkyl, alkoxy, alkylthio, alkylsulfinyl, alkylsulfonyl), were prepd. Thus, fuming nitric acid was added dropwise to 5-acetamido-3-bromo-1-(2,6-dichloro-4-trifiluoromethylphenyl)pyrazole and acetic anhydride in acetic acid; the mixt. was stirred at 60.degree. for 5 h to give 5-acetamido-3-bromo-1-(2,6-dichloro-4-trifiluoromethylphenyl)-4-nitropyrazole. Several I were effective against plutella xylostella larvae, all stages of Megoura viciae, and Spodoptera littoralis larvae.

IT 12015-83-59
RL: SPN (Synthetic preparation). PREP (Proceed)

12013-03-59
RL: SPN (Synthetic preparation); PREP (Preparation)
(prepn. of, as arthropodicide, nematocide, and anthelmintic)
12015-03-5 CAPLUS
HI-Pyrazole-3-carbonitrile, 5-amino-1-(2,4,6-trichlorophenyl)-4((trifluoromethyl)thio)- (9CI) (CA INDEX NAME)

L3 ANSWER 10 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 1993:191618 CAPLUS
DOCUMENT NUMBER: 118:191618
TITLE: Reactions of bromotrifluoromethane and related halides. Part 12. Transformation of disulfides into perfluoroalkyl sulfides in the presence of

sulfoxylate

anion radical precursors
Clavel, Jean Louis; Langlois, Bernard; Nantermet,
Roland; Tordeux, Marc; Wakselman, Claude
Rhone-Foulenc Rech., Cent. Rech. Carrieres,
Saint-Fons, 69192, Fr.
Journal of the Chemical Society, Perkin Transactions
1: Organic and Bio-Organic Chemistry (1972-1999)
(1992), (24), 3371-5
CODEN: JCPRB4; ISSN: 0300-922X
Journal
English
CASREACT 118:191618 AUTHOR (S):

CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

Perfluoroalkyl sulfides are prepd. by reaction of perfluoroalkyl halides with disulfides in the presence of sulfoxylate anion radical precursors. Aliph., arom. and heteroarom. disulfides bearing cyano, ester and amino functional groups have been employed; a variety of perhaloalkanes can

be employed, e.g., CF3(CF2)nI, CF3Br, CF2Br2, CF2BrC1, CFC13 and CF2C1CFC12. The most convenient sulfoxylate anion radical precursor for this reaction is formed by a combination of sodium formate and sulfur dioxide. Thus, reaction of pytazolyl disulfide I with HCC2Na and SO2 in DMF at 60.degree. and 12-13 bar for 4 h in an autoclave afforded perfluoroalkyl sulfide II in 85% yield. Also, reaction of PhSSPh with CF2BrC1 and Rongalite (sodium hydroxymethanesulfinate) in DMF-H2O at 1.7 bar and 20.degree. for 6 h afforded PhSCF2C1 in 72% yield. also

130735-50-9 (haloalkylation of, haloalkyl sulfide from) 130755-50-9 CAPLUS H-Pyrazole-3-carbonitrile, 4,4'-dithiobis(5-amino-1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

ANSWER 10 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

RN 130755-48-5 CAPLUS CN 1H-Pyrazole-3-carbonitrile, 5-amino-4-[bromodifloucomethy])thio]-1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

130755-51-0 CAPLUS 1H-Pyrazole-3-carbonitrile, 5-amino-4-[(bromochlorofluoromethyl)thio]-1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

146628-02-6 CAPLUS 1H-Pyrazole-3-carbonitrile, 5-amino-4-[(dichlorofluoromethyl)thio]-1-(2,4,6-tichlorophenyl)- (9CI) (CA IMDEX NAME)

Kamal Saeed

L3 ANSWER 10 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN

(Continued)

PAGE 1-A

PAGE 2-A

120115-83-5p 130755-48-5p 130755-51-0p 146628-02-6p

L3 ANSWER 11 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 1993:101949 CAPLUS
DOCUMENT NUMBER: 118:101949
TITLE: Preparation of
1-aryl-5-(arylalkylidentimnol pyrazoles
as pesticides
INVENTOR(S): Huang, Jamin: Ayad, Nafez Mohamed; Timmons, Philip
Reid
PATENT ASSIGNEE(S): Rhone-Poulenc Agrochimie, Fr.
SOURCE: EUR. Pat. Appl., 55 pp.
CODEN: EPXXDW
DOCUMENT TYPE: Patent

DOCUMENT TYPE: LANGUAGE:

English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

KIND DATE

Al 19921104 EP 1992-303857 19920425
B1 20011031 US 1991-693580 19910430
C1 19970827 RU 1992-802431 19920304
A 19930127 ZA 1992-802431 19920304
B6 20000216 C2 1992-3175 19920430
C1 US 1991-693580 A 19910430
US 1991-790449 A 19911112
US 1992-802431 A 19920304
US 1991-790449 A 19911112
US 1992-802431 A 19920304
CS 1992-1337 A 19920430 APPLICATION NO. DATE PATENT NO. KIND DATE EP 511845 EP 511845 EF 511845
R: AT, BE,
US 5236938
US 5360910
RU 2088576
ZA 9203175
CZ 286232 PRIORITY APPLN, INFO.:

OTHER SOURCE(S):

AB Title compds. [I; R1 = cyano, O2N, halo, CHO, (cyclo)alkylcarbonyl; R2 = halo, (halo)alkyl, (halo)alkoxy, NO2, SCN, (substituted) sulfamoyl, carbamoyl, alkoxycarbonyl, R8Son; R9 - (halo) (cyclo)alkyl, cycloalkylakyl, n = 0-2; R3 = H, alkyl, alkoxy, alkylthio, alkylamino; R4 = (substituted) heteroaryl, Ph; R5 = H, halo, alkyl; R6, R8 = H, F; R7 = (halo)alkyl; X = N, CR14 wherein R14 = H, halo, cyano, NO2, alkylamino; Qs = (halo)alkyl; X = N, CR14 wherein R14 = H, halo, cyano, NO2, alkyl alkylthio, alkoxyl, were prepd. Thus, a mixt. of

5-amino-1-(2,6-dichloro-4-trifluoromethylsulfenylpyrazole, 4-hydroxy-3-methoxybenzaldehyde, and p-toluenesulfonic acid were refluxed 40 h in PhMe with removal of H2O to give 85% title compd. II. Numerous I

II

L3 ANSWER 11 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued) as 100 ppm foliar or bait applications gave 70-100% kill of Spodoptera eridaria, Epilachna varivestis, Musca domestica, and Heliothis virescens.

17 14578-03-2P
R1: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIGL (Biological study); PREP (Preparation); USES (Uses) (prepn. of, as pesticide)
RN 145768-03-2 CAPLUS
CN 1H-Pyra2ole-3-carbonitrile,
1-(4-bromo-2,6-dichlorophenyl)-5-[[(4-bydroxy-3-methoxyphenyl)methylene]amino]-4-[(trifluoromethyl)thio]- (9CI) (CA INDEX NAME)

L3 ANSWER 12 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

Title compds. [I; X = halo, NOZ, (halo-substituted) alkylsulfenyl, alkylsulfinyl, alkylsulfonyl; Y = H, halo, cyano, alkylsulfenyl, alkylsulfinyl, alkylsulfonyl; Y = H, halo, cyano, alkylsulfenyl, alkylsulfinyl, alkylsulfonyl, alkoxy, (di)(alkyl)amino, trialkylammonium, cyanoalkylamino, alkoxyakylamino, alkoxyakylamino, alkoxyakylamino, alkoxyakylideneimino; Z = halo, cyano; RZ-RS = H, halo, (halo)alkyl, (halo)alkoxy, cyano, NOZ; gtoreq.l of RZ-RS H), were prepd. as insecticides, miticides, anthelmintics, protozoacides, and nematocides. Thus, NCGH:C(ONa)COZEt in ice water was treated with dil. RZSO4 to give crude ketoester, which was refluxed with Z-hydrazino-3-chloro-5-trifluoromethylpyridyl-3-ethoxycarbonyl-5-aminopycazole. This was condensed with C1FZCSC1 in AcON and the product was amidated with dimethylaluminum amide followed by dehydration with PCC13 to give title compds. II. Numerous I as 100 ppm foliar or bait applications gave 70-100% control of Aphis nasturtil, Spodoptera eridonia, Epilochna Varivestis, etc.
144292-76-2P 144292-89-7P
RL: SRN (Synthetic preparation); PREP (Preparation) (prepn. of, as arthropodicide, nematocide, anthelmintic, and protozoacide)
144292-76-2 CARLUS
1H-Pyrazole-3-carbonitrile, 5-amino-1-(3,5-dichloro-2-pyridinyl)-4-[(trifluoromethyl)thio]- (9CI) (CA INDEX NAME)

144292-89-7 CAPLUS
1H-Pyrazole-3-carbonitrile, 5-amino-1-(3,5-dichloro-2-pyridinyl)-4(methylthio)- (9CI) (CA INDEX NAME)

L3 ANSMER 12 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 1992:634010 CAPLUS
TITLE: 117:234010 Preparation of 1-(2-pyridyl)pyrazoles as pesticides
TITLE: 5 Phillips, Jennifer Lantz; Timmons, Philip Reid;
Powell, Gail Scotton; Pilato, Michael Thomas; Chou,
David Teh Wei; Huang, Jamin
Rhone-Poulenc Agrochimie, Fr.
SOURCE: EVERTON
DOCUMENT TYPE: CODEN: EPERTON
LANGUAGE: PRICE
English

FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

	A1 B1 CH, DE,	DK, ES, FR,	EP 1992-300467 GB, GR, IT, LI, LU	, MC, NL, PT, SE
R: AT, BE, NO 9200097 NO 179282 NO 179282 CA 2059088 CA 2059088 AU 9210251	CH, DE,	DK, ES, FR,	GB, GR, IT, LI, LU	, MC, NL, PT, SE
R: AT, BE, NO 9200097 NO 179282 NO 179282 CA 2059088 CA 2059088 AU 9210251	CH, DE,	DK, ES, FR,	GB, GR, IT, LI, LU	, MC, NL, PT, SE
NO 9200097 NO 179282 NO 179282 CA 2059088 CA 2059088 AU 9210251	70 '	100207220	NO 1002 07	10020108
NO 179282 NO 179282 CA 2059088 CA 2059088 AU 9210251	A B C AA C	19920720 19960603 19960911 19920719	NO 1992-97	19920108
NO 179282 NO 179282 CA 2059088 CA 2059088 AU 9210251 AU 644259 BR 9200219 IL 100678 CZ 281976 FI 9200221	B C AA C	19960603 19960911 19920719		
NO 179282 CA 2059088 CA 2059088 AU 9210251 AU 644259 BR 9200219 IL 100678 CZ 281976 FI 9200221	C AA C A1	19960911		
CA 2059088 CA 2059088 AU 9210251 AU 644259 BR 9200219 IL 100678 CZ 281976 FI 9200221	C A1	19920719		
CA 2059088 AU 9210251 AU 644259 BR 9200219 IL 100678 CZ 281976 FI 9200221	C Al		CA 1992-2059088	19920109
AU 9210251 AU 644259 BR 9200219 IL 100678 CZ 281976 FI 9200221	A1 .	20020618		
AU 644259 BR 9200219 IL 100678 CZ 281976 FI 9200221		19930128	AU 1992-10251	19920115
BR 9200219 IL 100678 CZ 281976 FI 9200221	B2 :	19931202		
IL 100678 CZ 281976 FI 9200221	Α :	19921006	BR 1992-219	19920116
CZ 281976 FI 9200221	A1 :	19960119	IL 1992-100678	19920116
FI 9200221	B6 :	19970416	CZ 1992-130	19920116
	Α :	19920719	FI 1992-221	19920117
JP 05086054	A2 :	19930406	JP 1992-44353	19920117
JP 3140829	B2 2	20010305	HU 1992-170 RO 1992-149204	
HU 62571	A2 :	19930528	HU 1992-170	19920117
HU 208534	В :	19931129		
RO 109940	B1 :	19950728	RO 1992-149204	19920117
PL 168/30	BI .	19960329	PL 1992-293228	19920117
RU 2088580	C1 :	19970827	RU 1992-5010813	19920117
CN 1063283	Α :	19920805	CN 1992-100330	19920118
CN 1041269	В :	19981223		
AT 158290	Ε .	19971015	AT 1992-300467	19920120
ES 2106821	тз :	19971116	ES 1992-300467	19920120
US 5306694	А.	19940426	CN 1992-100330 AT 1992-300467 ES 1992-300467 US 1993-79221	19930617
CN 1208036	Α :	19990217	CN 1998-106376	19980408
CN 1103771	в :	20030326		
ITY APPLN. INFO SOURCE(S):	. :	τ	S 1991-643530 A	10010110
SOURCE(S):			10 T217-042220 W	TAATOTIR

L3 ANSWER 12 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

L3 ANSWER 13 OF 15
ACCESSION NUMBER:
DOCUMENT NUMBER:
11991:429320 CAPLUS
115:29320
N-phenylpyrazole derivatives as insecticides

DOCUMENT TYPE: Patent English 1

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT N	ro.	KIND	DATE		APPLICATI		DATE
EP 41801	.6	A1	19910320				19900910
EP 41801	.6	B1	19950503				
R:	AT, BE,	CH, DE	, DK, ES,	FR, GB	, GR, IT,	LI, LU	NL, SE
ZA 90068	02	A	19911127		ZA 1990-€	802	19900827
NO 90039	08	A	19910312		NO 1990-3	908	19900907
AU 90623	112	A1	19910314		AU 1990-6	2312	19900907
AU 64923	10	B2	19940519				
CA 20249	55	AA	19910312		CA 1990-2	024955	19900910
HU 54868		A2	19910429		HU 1990-5	850	19900910
HU 20823	1	В	19930928				
CN 10532		A	19910724		CN 1990-1	07675	19900910
BR 90046	97	Α	19910910		BR 1990-4	697	19900910
DD 29764		A5	19920116		DD 1990-3	43914	19900910
RO 10725	5	В1	19931030		RO 1990-1	45905	19900910
PL 16364	12	B1	19940429		PL 1990-2	86822	19900910
AT 12203	18	E	19950515		AT 1990-3	09882	19900910
CZ 27947	6	В6	19950517		CZ 1990-4	387	19900910
ES 20717	177	T3	19950701		ES 1990-3	09882	19900910
JP 03118			19910520		JP 1990-2		19900911
JP 31000							
ORITY APPL					1989-2052	1 A	19890911
ER SOURCE			RPAT 115:	29320			

F3_mClmCS(0)n

L3 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER: 1991:5483 CAPLUS
DOCUMENT NUMBER: 114:5483
TITLE: Preparation of perhaloalkyl thioethers from disulfides

and perfluoroalkyl halides, and its application to pyrazole derivatives clavel, Jean Louis; Langlois, Bernard; Nantermet, Roland; Tordeux, Marc; Wakselman, Claude Rhone-Poulenc Agrochimie, Fr. EUR. Pat. Appl., 20 pp. CODEN: EFEXENW Patent French INVENTOR (S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PAT	ENT NO.	KIND	DATE		APPLICATION NO. DATE
FD	374061	A1	19900620		EP 1989-420489 19891212
	374061	B1			
22				. GI	R, IT, LI, LU, NL, SE
FR	2640264	A1	19900615	,	FR 1988-16710 19881213
	2640264	B1	19910125		
	2652810		19910412		FR 1989-13371 19891009
	2652810	B1	19930730		
	2004776	AA			CA 1989-2004776 19891206
	2004776		20000425		
	92639	A1	19961016		IL 1989-92639 19891211
	8906265	A	19900614		DK 1989-6265 19891212
UA	8946164		19900621		AU 1989-46164 19891212
	640621	В2	19930902		
HU	55738	A2	19910628		HU 1989-6508 19891212
HU	206661	В	19921228		
US	5082945	A	19920121		US 1989-448983 19891212
ES	2055145	Т3	19940816		ES 1989-420489 19891212
RU	2045517	C1	19951010		RU 1989-4742646 19891212
FI	95568	В	19951115		FI 1989-5938 19891212
FI	95568	С	19960226		
CZ	282729	В6	19970917		CZ 1989-7022 19891212
	1043499	A	19900704		CN 1989-109370 19891213
	1032201	В	19960703		
	02204477		19900814		JP 1989-323662 19891213
	2746707	B2	19980506		
	8906521	A			BR 1989-6521 19891213
	8909519	A	19910828		ZA 1989-9519 19891213
	5283337	A	19940201		US 1991-789332 19911108
RITY	APPLN. IN	0.:			1988-16710 A 19881213
					1989-13371 A 19891009
			nnn 114.540		1989-448983 A3 19891212

MARPAT 114:5483 OTHER SOURCE(S):

L3 ANSWER 13 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

The title compds. (I; A = iodo, Br, H, NH2; m = 1,2; n = 0, 1, 2), useful for controlling arthropod, plant nematode, helminth, or protozoal pests, are prepd. Thus, a soln. of I [A = NH2, F3-mclmcs(O)n = CHClF2S] in dry THF was added to tert-BuONO2 at room temp, and the mixt was stirred 3 days at room temp, to give I [A = H, F3-mclmcs(O)n = CHClF2S]. I at .ltoreq.500 ppm gave 60% mortality against the larvae of Plutella xylostella. 120115-83-52

120115-83-59
RK: SPN (Synthetic preparation); PREP (Preparation)
(prepm. of, as intermediate for pesticidal phenylpyrazole)
120115-83-5 CAPUS
HR-Pyrazole-3-carbonitrile, 5-amino-1-(2,4,6-trichlorophenyl)-4[(trifluoromethyl)thio]- (9CI) (CA INDEX NAME)

L3 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

AB Perhaloalkyl thioethers are prepd. by reaction of disulfides with perfluoroalkyl halides and reducing agents formed from (a) SO2 and either Zn, Cd, Al, or Mn, or (b) an alkali metal dithionite, or (c) an alkali metal, alk. earth, or other metal hydroxymethanesulfinate, or (d) a formate and SO2. For example, reaction of Ph2S2 with Na dithionite and CF3Br(g) in aq. DMF conts, Na2HPO4 at 20. degree. gave 65 k PhSCF3. Pyrazole deriv. I was similarly prepd. using SO2 and Na formate with 95% conversion and 90% yeld. Various aliph., arom., and pyrazole-derived thioethers were prepd; yields ranged from 6 to 93%.

II 130755-49-6F
RI: RCT (Reactant); SPN (Synthetic preparation); PREF (Preparation); RACT (Reactant or reagent)
(prepn. and hydrolysis-dimerization of)
RN 130755-49-6 CAPLUS
Thiocoyanic acid,
5-amino-3-cyano-1-(2,4,6-trichlorophenyl)-1H-pyrazol-4-yl ester (9CI) (CA INDEX NAME)

130755~50-9P 130755-50-99
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) [prepn. and reaction of, with fluoroalkyl halides and reducing agents) 130755-50-9 CAPLUS
H-Pyrazole-3-carbonitrile, 4,4'-dithiobis[5-amino-1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

L3 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN

(Continued)

PAGE 1-A

PAGE 2-A

IT 130755-48-5P 130755-51-0P
RL: SPN (Synthetic preparation); PREP (Preparation)
(prepn. of, by reaction of disulfide with fluoroalkyl halide and reducing agent)
RN 130755-48-5 CAPLUS
CN 1H-Pyrazole-3-carbonitrile,
5-amino-4-[(bromodifluoromethyl)thio]-1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

L3 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

L3 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN

Br-CF2-S NH2

130755-51-0 CAPLUS
1H-Pyrazole-3-carbonitrile, 5-amino-4-[(bromochlorofluoromethyl)thio]-1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

(Continued)

T 130755-52-1P 131960-95-7P
RL: SPN (Synthetic preparation); PREP (Preparation)
(prepn. of, via reaction of disulfide with reducing agent and fluoroalkyl halide)
N 130755-52-1 CAPLUS
N 1H-Pyrazole-3-carbonitrile,
-amino-4-[[bromochlorofluoromethyl]sulfonyl]1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

RN 131960-95-7 CAPLUS
CN 1H-Pyrazole-3-carbonitrile,
5-amino-4-(bromochlorofluoromethyl)sulfinyl]1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

L3 ANSWER 15 OF 15
ACCESSION NUMBER: 1990:35845 CAPLUS
DOCUMENT NUMBER: 112:35845
TITLE: 8
Plants, N-phenylpyrazole derivatives as pesticides for

animals, and man, and their preparation,

compositions,

and use
Buntain, Ian George; Hatton, Leslie Roy; Hawkins,
David William; Pearson, Christopher John; Roberts,
David Alan
May and Baker Ltd., UK
Eur. Pat. Appl., 40 pp.
CODEN: EPXXDW
Patent
English
4 INVENTOR (S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT:

PAT	TENT NO.		KIND	DATE		A		CATI	ои ис	٥.	DATE
	295117		A1	19881214		E			0530	5	1988061
EP	295117		B1	20000405							
		BE,	CH, DE	, ES, FR,							
	86492		A1	19930708					6492		1988052
	105138		A1	19940826					0513		1988052
	8803140		A A	19881213		D	19	88-3	140 735		1988060
	8802735		A	19881213							1988060
	8802551		A	19881213		ИС	19	88-2	551		1988060
	175367		B C	19940627							
	175367										
	8817554		A1	19881215		AU	19	88-1	7554		1988060
	618266		B2	19911219							
	100612		B1	19920707					33912		1988060
	106496		B1	19930531					4435		1988060
	63316771		A2	19881226					4345		1988061
	8804179		A	19890222					179		1988061
	48875		A2	19890728		н	19	88-3	009		1988061
	203729		В	19910930							
	153478		B1	19910430					72998		1988061
	1330089		A1	19940607		C.F	19	88-5	69272	2	1988061
	210668		В	19950628		HU	19	91-1	577		1988061
	278972		В6	19980506		51	19	88-4	052		1988061
	285151		В6	19990512		Cz	19	88-4	052		1988061
EP	967206		A1	19991229					1379		1988061
		BE,	CH, DE	, ES, FR,	GB,						SE
	191479		E	20000415					0530		1988061
	2144390		T3	20000616		ES	19	88-3	0530	5	1988061
	B8103601		A	19881228		Ch	19	88-1	0360	ι	1988061
	1027341		В	19950111							
	9701475		B1	19970206		KF	19	88-7	045		1988061
	8803258		A	19890131		BF	19	88-3	258		1988061
	281744		A5	19900822		DI	19	88-3	16723	3	1988061
	281744		В5	19970220							
	2051909		C1	19960110					89476		1991031
	9501839		A	1995041B		F1	19	95-1	839		1995041
	1005289		A1	20010209		HE	19	98-1	02258		1998031
	3033663		T 3	20001031					01350		2000061
D.12	20020152	7	A5	20021010		D.M	- 20	02-1	527		20021010

L3 ANSWER 15 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN IL 1988-86492 DK 1988-3140 FI 1988-2735 EP 1988-2735 (Continued)
A 19880525
L 19880609
A 19880609
A3 19880610
A 19880610 HU 1988-3009

OTHER SOURCE(S):

MARPAT 112:35845

The title compds. [I; R1 = cyano, NO2, halo, Ac, CHO; R2 = R5S(O)n where

The title compds. [I; R1 = cyano, NOZ, halo, Ac, CHO; R2 = R58(0)n where

= 0, 1, or 2; R5 = (.ltoreq.1 halo-substituted) straight- or
branched-chain .gtoreq.4 alkyl, alkenyl, or alkynyl; R3 = H, NR6R7, halo,
straight- or branched-chain C2-5 alkoxymethyleneamino (un)substituted on
methylene by a straight- or branched-chain C1-4 alkyl; R6, R7 = H,
straight- or branched-chain .ltoreq.5 alkyl, alkenylalkyl, or
alkynylalkyl, CHO, (.ltoreq.1 halo-substituted) straight- or
branched-chain C2-5 alkanoyl or alkoxycarbonyl, or NR6R7 = 5- or
6-membered cyclic imido: R4 = 2- or 6-halo- or 4-straight- or
branched-chain (C1- or Br-substituted) alkyl- or alkoxy-substituted
phenyl; with the exclusion of the compd. wherein R1 = cyano, R2 = MeSOZ,
R3 = NH2 and R4 = 2,6,4-C12(CF3)C6H2], useful for control of arthropod,
plant nematode, helminth and protozoan pests (no data except insects),
were prepd. A stirred soln. of 20 g 5-amino-3-cyano-1-(2,6-dichloro-4trifluoromethylphenyl)pyrazole in CH2C12 was treated dropwise with a

soln.

of 10.8 g CF3SC1 in CH2C12 during 1 h. The resulting soln. was stirred overnight at room temp. to give 24.2 g 5-amino-3-cyano-1-(2,6-dichloro-4-trifluoromethylphenyl)-4-trifluoromethylthiopyrazole (II). I at <500 ppm caused at least 65% mortality against Plutella xylostella larvae. A water-sol. conc. was formulated from II 7, Ethylan BCP 10% w/v and N-methylpyrrolidone 1004 by vol.

IT 120069-19-4

RL: RCT (Reactant); RACT (Reactant or reagent) (xxidn. of)
RN 120069-19-4 CAPLUS
CN 1H-Pyrazole-3-carbonitrile,
5-amino-4-[(1-methyl-2-propynyl)thio]-1-(2,4,6-trichlorophenyl)- (9CI) (CA INDEX NAME)

L3 ANSWER 15 OF 15 CAPLUS COPYRIGHT 2004 ACS on STN (Continued)

TT 120115-83-5P

120115-83-5P
RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of, as pesticide)
120115-83-5 CAPLUS
11-Pysazole-3-carbonitrile, 5-amino-1-(2,4,6-trichlorophenyl)-4-((trifluoromethyl)thio]- (9CI) (CA INDEX NAME)

=> logoff

ALL L# QUERIES AND ANSWER SETS ARE DELETED AT LOGOFF

LOGOFF? (Y)/N/HOLD:Y

COST IN U.S. DOLLARS

SINCE FILE TOTAL
ENTRY SESSION
72.22 227.85

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL

CA SUBSCRIBER PRICE ENTRY SESSION -10.40 -10.40

STN INTERNATIONAL LOGOFF AT 15:33:08 ON 10 JUN 2004